REMARKS

This paper is responsive of an Office Action mailed on September 20, 2005. Prior to this response claims 1-13 and 27-28 were pending. Claims 1-13 and 27-28 remain pending.

Claims 1-13 and 27-28 have been rejected under 35 U.S.C. 102(e) as anticipated by Huotari (US Pub 2004/0106249). The Office Action states that Huotari describes all the limitations of claims 1, 27, and 28. This rejection is traversed as follows.

The priority date for the Huotari application is December 3, 2002, based upon Provisional Application 60/430,960. To "swear behind" Huotari's priority data, the Applicant has enclosed, as Attachment A, the declaration of Wei Goa, one of the co-inventors of the instant application. In his declaration, Mr Goa swears that the subject matter of the instant application claims were conceived of, and reduced to practice prior to the date of December 3, 2002.

Attachments B and C are enclosed to support Mr. Goa's assertions. Attachment B is a true copy of the patent disclosure that was submitted to the Sharp Laboratories of America (SLA) Patent Department. The instant application was authorized as a result of the patent disclosure. The patent disclosure includes a figure on page 4 of a gate electrode comprising a gate dielectric, a barrier metal overlying the dielectric, and a gate metal overlying the barrier metal. Some examples of barrier metals are mentioned in Section 10(2). Gate metals are discussed in Section 10(3). Section 10(4), under the figure, states that the thinness of the barrier metal precludes it from having an affect on the gate electrode work function.

Attachment C is a true copy of a monthly report submitted by Wei Goa to his supervisor (and co-inventor) Yoshi Ono. Table 2 (incorrectly

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marked as "Table 1") and the accompanying description discuss the measured work function of various gate electrode stacks (gate metal/barrier layer). For example, shown in the Table are Pt/TaN and Nb/TaN stacks. Further, page 1 of the report discusses the work function of Pt/TiN and Ir/TiN stacks. The monthly report clearly shows that the claimed invention was reduced to practice. In his declaration, Mr. Goa swears that this reduction to practice occurred prior to December 3, 2002.

Since claims 1-13 and 27-28 were conceived of, and reduced to practice prior to the priority date of the prior art reference, the Application respectfully requests that the rejection be withdrawn.

It is believed that the application is in condition for allowance and reconsideration is earnestly solicited.

Date: 12/19/2005

Gerald Maliszewski Registration No. 38,054

Respectfully submitted,

Customer Number 55,286

P.O. Box 270829

San Diego, CA 92198-2829

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gerry@ipatentit.net

ATTACHMENT A

Patent Application Serial No. 10/784,662 Filed February 23, 2004

DECLARATION OF WEI GAO UNDER 37 CFR 1.131

- I, Wei Gao, hereby declare as follows:
- 1. My residence address is 2402 SE 180th CT, Vancouver. WA 98683.
- 2. Since April 2001, I have been employed by Sharp Laboratories of America, Inc. (SLA), 5750 N.W. Pacific Rim Blvd., Camas, WA 98607.
- 3. I am a co-inventor of Patent Application Serial No. 10/784,662, and I invented the device recited in amended claims 1-13 and 27-28 prior to December 3, 2002.
- Attachment B is a true copy of the SLA patent disclosure document filed with the SLA Patent Department. I affirm that the patent disclosure document was written, signed, and witnessed before December 3, 2002. The disclosure shows the concept of a gate electrode having a first metal barrier layer, with an overlying second metal, where the first metal layer is thin enough so as to not effect the overall work function of the gate electrode.
- 5. Attachment C is a true copy of weekly report describing the results of experiments conducted in the laboratory. The report shows a first metal buffer layer of TiN, TaN, or Ti, having a thickness in the range of 1 to 200 nanometers (nm). The overlying second metal layer is Al, Nb, or Pt. The lab results and write clearly show a reduction to practice of the claimed invention.
 - I hereby declare that all statements made herein of my

own knowledge are true and that all statements made on information and belief are believed to be true, and further that these statements were made with the knowledge that willful, false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United State Code and that such willful, false statements may jeopardize the validity of the application or patent issuing thereon.

12/5/2005

Date

Wei Gao

ATTACHMENT B

PAGE 12/22 * RCVD AT 12/19/2005 9:44:51 PM [Eastern Standard Time] * SVR:USPTO-EFXRF-6/26 * DNIS:2738300 * CSID:8584519869 * DURATION (mm-ss):05-38

Invention Disclosure

Inventor Signature

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PAT-DIS_Metal Gate1.doc

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SHARP Laboratories of America, Inc.	SLA Confidential	Page 2 of 5
5750 NW Pacific Rim Boutevard. Camas, Washington 98607	SLA Docket No.	674
Names	Sheng Teng Victor Hsu	
Supervisor's Name:	Director, IC Process Technology	
Supervisor's Title:		
Project Number/Name:		
4. Conception of the Invention:		
Date Conceived:		
Date of first Written Description:	Notes for Quarterly Rev	iew Meeting
Notebook & Page No. or File Archive:	Notes (c) deal	
Date first explained to others (whom?):		
Planned Application for the Invention:		
6 Construction & Test of First Prototype I	Embodying the Invention:	
Date First Prototype Completed:		
erant gunnessful Test:		
Successful Operation Witnessed By:		
6. Public Disclosure of Invention (Presen (NOTE: Patent Application MUST be fil		eation) .):
Date of Luce -		
Setting (Conference/Journal Name):		
Title of Paper or Presentation:		
Time of Displacure (Written/Verbal):		
Does Data Sheet or Application Note Disclose	e the Invention (when)?	
7. What is the field of the invention (Invention	ention relates to):	
This invention relates to integrated ci characteristics.		
8. What is the problem solved by your i search pages here)?		
Doped polysilicon has been the gas of microelectronics technology. To performance), p+ poly is used for scaled, the thickness of the polystresistance and a targe effective or	PMOS and n+ poly is used for N	IMOS. As devices are maintain low sheet
This sand		
Inventor Signature Date	- 06 · 08	
The state of the s	Witnessed & Understood By	0 05
Invertor Signaturo	3	Date*
Inventor Signature Date	Winessed & Understood By	
	vention Disclosure	PAT-DIS_Metal Gate1.do

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Camas, Washington 98607

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Page 3 of 5

SLA Docket No.

674

has been necessary to increase poly doping density with each successive generation. This has led to the problem of channel autodoping in which B from the gate poly diffuses across the thin gate dielectric and into the channel, causing Vth variations which degrade device performance.

In order to solve the problems of gate resistance, poly-depletion, and B-diffusion, the industry soon will be forced to switch to metal gate electrodes. To achieve acceptable Vth, metal gate materials with the correct work function will be necessary. This implies one metal for NMOS and a different metal for PMOS. Thus, dual metal (with work functions corresponding to p+ and n+ poly Si) will be required for CMOS circuits.

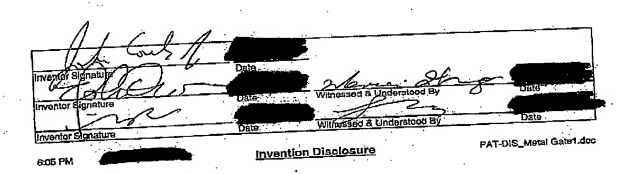
It has been shown that many of the desirable metal materials have adhesion and/or stability problems when placed in direct contact with the SiO_2 or high k gate dielectrics (HfO₂, ZrO₂, etc.). For example, it is known that Pt does not adhere well to SiO_2 and metals such as Ti, Hf, Zr, etc. scavenge O, reducing the underlying dielectric film, causing degradation and increased leakage.

This method will enable the use of certain metal and dielectric combinations for the manufacture of integrated circuits.

9. How is your solution different from the prior art (one paragraph or list)?

Metal gates will be needed in future MOS devices. In previous work, metals have been placed in direct contact with the gate dielectric (SiO2 or high-k) and revealed adhesion and stability issues (reactivity of metal with dielectric leading to oxygen scavenging)

We disclose the use of a thin Interfacial layer / barrier to 1) improve adhesion between metal gate and dielectric and / or 2) serve as a barrier metal to reduce reactivity. A key element is that, based on an earlier SLA disclosure (W. Gao and Y. Ono), as long as the thickness of this interfacial metal is < ~50A, it will not impact Vth of the transistor. Because the barrier / layer is a metal, gate resistivity will not be adversely impacted.



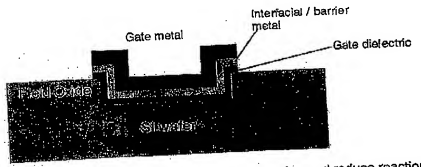
PAGE 14/22 * RCVD AT 12/19/2005 9:44:51 PM [Eastern Standard Time] * SVR:USPTO-EFXRF-6/26 * DNIS:2738300 * CSID:8584519869 * DURATION (mm-ss):05-38

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SLA Confidephal	Page 4 of 5
SLA Dacket No.	

- 10. Please give a detailed description of your invention, include any graphics, notebook pages or other material necessary to understand your invention.
 - (1) The structure prior to depositing the barrier layer is formed by any state of the art method. The example shown in the figure is for the gate replacement process. A gate dielectric of any kind has been deposited / grown and is ready for the metal gate.
 - (2) A thin barrier / interfacial layer of metal of desired thickness (< 50A) is deposited by any state of the art method. Depending on the structure, this metal material can be TiN, TaN or WN.
 - (3) A metal gate with the desired workfunction / properties is then deposited to desired thickness by any state of the art method. Depending of the design of the structure, this film can be Pt, Ti, Hf, Zr, or any other potential metal or metal compound.
 - (4) The metal stack is then etched or CMP'd to form the device structure desired. The structure can then be treated thermally, electrically, or mechanically, as required.

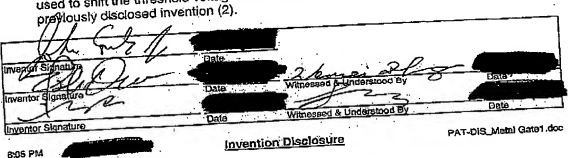


The barrier / interfacial metal should improve adhesion and reduce reaction between the overlying gate and the underlying dielectric. Because the interfacial layer is <-50A, the workfunction of the resulting device should be determined solely by the top gate metal.

11. What other embodiments or examples are there of your invention?

We envision that potentially, a gate stack consisting of any gate metal and any dielectric could be made compatible, providing a suitable barrier metal can be found.

In another embodiment, should the workfunction of the metal gate not be an optimum value, the proper thickness of the intermediate layer, typically greater than 90Å can be used to shift the threshold voltage to the desired value. This is in accordance to the



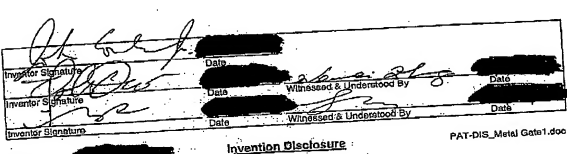
PAGE 15/22 * RCVD AT 12/19/2005 9:44:51 PM [Eastern Standard Time] * SVR:USPTO-EFXRF-6/26 * DNIS:2738300 * CSID:8584519869 * DURATION (mm-ss):05-38

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SLA Confidental	Page 5 of 5
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SLA Docket No.	474

- List related publications, patents, articles, or other references and give a brief summary (Indicate the extent of your search, attach search pages, if any):
 - 1. V. Misra, G.P. Heuss, H. Zhong, "Use of metal-oxide-semiconductor capacitors to detect Interactions of Hf and Zr gate electrodes with SiO₂ and ZrO₂," Appl. Phys. Lett. 78,
 - 2. W. Gao and Y. Ono, "MOSFET threshold voltage tuning with metal gate stack control," SLA Patent Disclosure,
 - 3. V. Misra, M. Kulkarni, G. Heuss, H. Zhong, and H. Lazar, "Electrical and Material Properties of Metal Silicate Dielectrics and Metal Gate Electrodes for Advanced CMOS Devices," in Electrochemical Society Proceedings Vol. 2000-9, p. 291-98 (2000).



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ATTACHMENT C

Sharp Labs of America

To:

Yoshi Ono

From:

Wei Gao

cc:

Group Files

Date:

Subject:

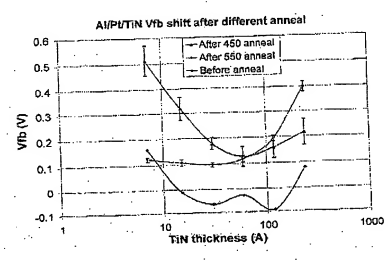
Single metal gate lot 3

Single metal gate lot 3 has gone through radical gate oxidation and RTA. The detailed information is listed in table 1 and the lot is at contact etch and is going to be ready for measurement soon.

	le 1. Radical oxidation	parameter and F	TA anneal
l an	Wafer temperature	Time(min)	Wieastifed Oxide attendess
Radical oxidation	200°C	8	29A 27Å
RTA(MPT)	750°C	1	2/12

Dual metal gate project

PVD TaN using 50%N₂/Ar produced in our lab has show its stability with SiO₂ up to high temperature. However, the workfunction of this material is not stable even after 450°C anneal. From workfunction tuning point of view, Pt/TiN or Ir/TiN should provide big enough V_{tb} shift although the work function does not lineup with Si bandgap very well. However, our TiN is known to be oxidized at elevated temperature, and we don't have good enough furnace to anneal known to be oxidized at elevated temperature, and we don't have good enough furnace to anneal the device without oxidizing them. In order to try our TiN stack with Pt, we put 2000Å Al on top of Pt(1200Å)/TiN to see the effect of workfunction tuning before/after anneal, the result is showing in Figure 1.



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Figure 1 shows that before anneal, Al/Pt/TiN stack shows a scattered result. Not only the curve goes up and down, the error bar is also too big to put on the graph. After 450°C anneal, V_{tb} shows an smooth change up to 60Å and goes the other way after that. The V_{tb} change is around 0.4V. After 550°C anneal, the curve further changed which could be attributed to the involvement of Al since Al could diffuse through Pt. On the left, (7Å point), work function of Pt seems greatly dropped (because of the Al) and on the other side, the work function of 236Å TiN increased for some other unknown reason. Besides all this dramatic changes, the Cacc is observed to have certain change but the capacitor doesn't fail and the change is limited.

So far we have tested several workfunction tuning material combinations based on our dual metal gate configuration. Table 2 summarize the result we got so far.

dual literar San	_			·	nte
	Table 1. S	ummary of wo	rkfunction tul	ning experime	Possible reason
stack	Transition	V _{fb} shift (V) as deposited	After 450°C anneal (v)	Cacc	
	region	0.6	0.5	Not stable	Al/SiO ₂
AI/TiN	100 Å	0.9	0.9	Not stable	Ti/SiO ₂
Pt/Ti	40 Å	N/A	0.25	Not stable	Ta/SiO ₂
Pt/TaN*	100 Å	N/A	0.3	Not stable	Ta/SiO ₂ and Al
Al/TaN*	100 Å	0.75	N/A	Stable	TaN/SiO ₂ is stable
Al/TaN**	50 Å	0.8	0.3	Stable	workfunction of TaN
Nb/TaN**	100 Å	0.8	0.2	Stable	is not stable
Al/Nb/TaN** Al/Pt/TiN	100 Å 60 Å	N/A_	0.4	Stable	Al involvement
AUPUID	1 YV				

^{*} Ta rich PVD TaN using 15%N2/Ar

From this table we can see the first 4 experiment proves that the dual metal gate stack require both metal to be stable when contact with SiO2. and the next 4 experiment shows that the stack is stable with SiO2, however, the workfunction properties of the stack is not stable during anneal. As good barrier layer or adhesion enhancer, TaN and TiN all have multiple valence states which allows them to form many nitrides with different stoichiometries, PVD, on the other hand, generates meta-stable film structure which allows flexibility to change easily during anneal.

Besides the stability issue, the sputtering process always causes gate dielectric thinning as well as introducing charges, the loss becomes bigger when the gate dielectric is getting very thin and the extra charge introduced may require unnecessary treatment than there is not. Generally speaking, CVD or ALD is more preferred process when work with gate dielectric.

We have decided to try CVD TaN on our CVC system and literature search was carried out. Since there are only a few published research on CVD or ALD TaN, I listed the major feature of different approaches into the following table.

From the details listed in the table, the following conclusion could be drawn:

- 1. From barrier properties point of view, CVD or ALD TaN is preferable. However, the film densities are normally lower (<10) which require post deposition anneal for densification and stabilization. PVD TaN has higher density (>10) but still lower than
- 2. As a workfunction tuning material, it has to be stable and high density, as deposition method, the substrate temperature has to be lower than 430°C (to fit in CVC system and not to interfere with our existing process). Judged by this two categories, TBTDET ALD is no good because the deposition temperature has to be <260°C and the film has low density. PEALD TBTDET can produce higher density film under 260°C, however, it

^{**} N rich PVD TaN using 50%N2/Ar

- requires modification of our CVC tool (add plasma system). Y. H. Kim et al used CVD TBTDET to produce TaN that is stable up to 1000°C, however, the deposition temperature is 600°C followed by 700°C anneal in NH₃.
- 3. Tantalum halide (Cl and Br) were studied to deposit TaN as the candidate of Cu barrier layer three years ago, due to the bonding strength difference, TaCl₅ require >600°C deposition temperature in order to get low resistivity, while TaBr₅ require only 350-425°C. The film deposited from CVD TaBr₅ contains <2% O and <0.5%Br, the film is amorphous TaN_{1.83} and its resistivity is normally higher.
- 4. Because of the constrain on deposition temperature, CVD TaBr₅ becomes the only approach that could fit in to our current CVC system.
- 5. It seem CVD TaBr₅ could produce so far the purest TaN at low temperature compare with other ALD or CVD methods. As a Cu barrier material it fails at above 550°C due to the fast boundary diffusion when the film turns from amorphous to microcrystalline. For our dual metal gate project, this may not be a problem. The major questions for TaBr₅ produced TaN are its resistivity, its workfunction, and workfunction thermal stability.

		C ASCCome	nt deposition	approaches	for TaN film	1
Table 2.	Comparison	CVD	ent deposition	CVD		
	CVD TaBr ₅	TaCl ₅	TBTDET	TBIDET	TBTDET	sputtering 300°C
Deposition temperature	350-425°C	>600° C	260°C	600°C d. 700°C a.	<260°C[1]	300 C
ΔH (kcal/mol)	-143	-205	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \		NH ₃	
Reducing agent Crystal phase	$H_2 + NH_3$ $TaN_{1.83}[2]$		H ₂ Poly	TaN [3]	amorphous	TaN _{1.71} [2] poly
Density	amorphous 9.7[2]		7.9[1]		3.6[1]	~13[0], 10.17[2]
Contamination	<2%O		15-35%C and O	1.5%C, <1%O[3]	Significant amount	
ρ(μΩcm)	2500		400			~ 400
Thermal stability	< 550°C			1000°C		<650°C[2
Step coverage	90% at 8:1		100% at			Poor
Major drawbacks	V.15µm	High temp.	Slow proc	High temp.	Low den. aging eff.	Step coverage

^[0] Our PVD data.

^[1] J-S Park, H-S Park and S-W Kang, J Elec. Chem. Soc. 149(2002)C28

^[2] A. E. Kaloyeros et al. J Elec. Chem. Soc. 146(1999)170

^[3] Y. H. Kim, et al., IEDM 2001

ATTACHMENT D

DEC 1 9 2005

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:

Attorney Docket: SLA0674

Inventor:

John F. Conley, Jr., et al.

Confirmation No. 2287

Serial No.:

10/784,662

Group #: 2815

Filed:

February 23, 2004

Examiner: Edward J. Wojciechowicz

Title:

REACTIVE GATE ELECTRODE CONDUCTIVE BARRIER

Commissioner for Patents P.O. Box 1450 Alexandria, Virginia 22313-1450

ASSOCIATE POWER OF ATTORNEY

CHANGE OF CORRESPONDENCE ADDRESS

I hereby appoint Gerald Maliszewski, P. O. Box 270829 San Diego, CA 92198-29, U. S. Patent and Trademark Office Registration Number 38,054, as an associate attorney for the above-captioned matter, with full power of substitution and revocation, to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith.

Respectfull

Please change the Correspondence address to that associated with Customer Number 55,286.

David C. Rimma, Registration # 27,6

Data

David C. Ripma Patent Counsel

Sharp Laboratories of America, Inc.

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Camas, WA 98607

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Page 1 Associate Power of Attorney for Serial No. 10/784,662